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Exhibit "K" attached to Declaration of John C. Rockett. Ph.D.

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Macroresults through Microarrays

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The third enactment of Cambridge Healthtech Institute's Macroresults through Microarrays meeting was held in Boston (MA, USA) from 29 April-1 May 2002. The subtheme of this year's meeting was 'advancing drug discovery', a widely touted application for array technology.

The evolution of microarrays

If you were asked 'Who first conceived of the idea of microarrays', who would come to mind? Mark Schena perhaps, first author of the seminal 1995 paper on cDNA arrays [1]? Maybe Pat Brown, Schena's then supervisor? Or perhaps Stephen Fodor, the primary driver behind Affymetrix's (http://www. affymetrix.com) oligonucleotide-based platform [2]. Brits might even chant the name of Ed Southern [3]. Well, according to Roger Ekins (University College London Medical School; http://www. ucl.ac.uk/medicine/) all these answers would be wrong. It was in fact Ekins and his colleagues who first conceivedof and patented 'a new generation of ultrasensitive, miniaturized assays for protein and DNA-RNA measurement based on the use of microarrays' in the mid 1980s [4]. The concept and potential of array technology was more fully described in a later publication, in which Ekins et al. [5] concluded that antibody microspots of -50 μm² could be achieved, and that as many as 2 million different immunoassays could, in principle, be accommodated on a surface area of 1 cm².

Technological innovation

In practice, it took a different biological molecule (DNA), a different research

group, and a leap into microfabrication technology to even begin approaching these kinds of densities [Affymetrix patent 6045996 talks of one million spots cm-2]. Of course, advancing technology is one of the driving engines behind the genomics juggernaut, and we are already seeing '4th generation' machines for fabricating DNA chips. If the company representatives at this meeting are to be believed (and their cases seemed strong), spotting is out, and in situ fabrication of oligonucleotide-based 'iterative custom arrays' is in. Whether you go with the Combimatrix's (http:// www.combimatrix.com) electrochemically directed synthesis and detection system, febit's (http://www.febit.com) Geniom® technology, or Nimblegen's (http://www.nimblegen.com) Maskless Array Synthesizer technology is a matter of personal choice. However, each of these machines provides the flexibility to design variable length oligonucleotide probes from sequences inputted by the user, and then perform in situ synthesis of an array. Each system also boasts unique advantages. For example, Combimatrix's biological array processor is a semiconductor coated with a 3D layer of porous material in which DNA, RNA, peptides or small molecules can be synthesized or immobilized within discrete test sites, while febit's Geniom One® is a fully integrated gene-expression analysis system with minimal user hands-on time - the probe sequences are programmed, the RNA samples inserted, and the gene expression data is pumped out a few hours later.

Cell- and tissue-based arrays

Array technology is in most people's minds firmly linked with gene-expression profiling. Fewer are aware that cell- and tissue-based arrays have been developed, and how they can provide a vital extra dimension to research. In support of this, Barry Bochner gave an update on the cell-based array system that Biolog (http://www.biolog.com) has produced for simultaneously measuring the effects of one gene in the cell under thousands of growth conditions (see [6] for further details). David Walt (Tufts University; http://www.tufts. edu/) is developing single live cell arrays using optical imaging fiber (OIF) technology. An array of microwells is fabricated on the face of an OIF at densities of up to 10 million wells cm-2. Cells are then added to the wells and disperse at an average of one cell per well. Physiological and genetic responses of each cell are measured via fluorescence produced by reporter genes (e.g. lacZ, gfp. Assays performed so far include yeast live or dead cell assay, microenvironment pH and O₂ measurements, promotor responses using the lacZ and phoA reporter genes, and protein-protein interactions using the yeast two-hybrid system. The main advantage of this system is that the cells remain alive during the assay, which means a real-time timecourse can be performed and/or the array passed from sample to sample. This would be useful in, for example, the scanning of a combinatorial drug library for specific physiological effects.

Tissue arrays are a useful complementary technology to DNA arrays because they can be used to help validate and

understand the biological and medical significance of gene changes discovered using standard DNA arrays. For example, an array of tumor tissues can be screened for the protein (using immunohistochemistry), message (using in situ hybridization) and copy number (using comparative genomic hybridization) of a gene of interest, to determine if expression of the gene (or lack thereof) is related in any way to survival. They can also be used to predict the probability of clinical failure of lead compounds as a result of toxicity by evaluating the distribution of the drug targets in normal tissue. Spyro Mousses and his co-workers at the National Human Genome Research Institute (http://www.nhgri.nih.gov/index.html) have built such arrays, including a multi-tumor array (-5000 specimens, and sections from 36 normal and 800 metastatic tissues) and a normal tissue array (76 tissue and 332 cell types).

The problem with proteins

It has been said that genomics tells us what might happen, transcriptomics indicates what should happen, and proteomics shows what is happening. The impact of functional proteomics on pharmaceutical R&D is rapidly increasing, and protein arrays are being used increasingly in both basic and applied research. Their use lies not only in comparative protein expression and interaction profiling, but also in diagnostics and drug discovery. However, an increasing number of researchers have found that protein arrays, like their cousins the DNA arrays, present several practical obstacles relating to their production and use. For example, in using Escherichia coli to produce recombinant eukaryotic proteins from a single expression vector, multiple protein products are often produced, suggesting mixes of truncated or otherwise altered proteins. There is also the obvious concern that the proteins might not be modified in a similar manner to

eukaryotic systems. Also, an optimal method for depositing and binding proteins to the selected substrate is yet to be determined, as is the best way to ensure that they are bound in a correctly folded, active conformation.

Several companies have been addressing these problems. Prolinx (http:// www.prolinxinc.com) is one such company, and Karin Hughes described their Versalinx™ chemistry for producing protein, peptide and small-molecule arrays. Versalinx™ uses solution-phase conjugation followed by immobilization, resulting in functional orientation of proteins and peptides on the substrate surface. It also offers the valuable additional benefit of exhibiting low non-specific binding. Sense Proteomic (http://www.senseproteomic.com) is also among those addressing these problems to develop robust protein arrays for drug discovery and clinical applications and has developed functional protein array formats based on specific disease tissues. Subtractive hybridization is used to identify genes with altered expression in breast tumor and cystic fibrosis compared to normal tissue. A high throughput cloning strategy (COVET™) is then used to produce libraries of genes that are tagged, cloned, expressed, purified and finally immobilized on glass slides. Initial validation studies have shown that the vast majority of the immobilized proteins do indeed retain biological function.

Stefan Schmidt and his company (GPC Biotech; http://www.gpcbiotech. de) have moved past the platform development stage and, with their focus firmly on drug discovery, are currently developing kinase-profiling arrays. Kinases are important targets for pharmaceutical drug discovery and therapy, and GPC's aim is to simultaneously detect multiple kinases, obtain activity profiles for different cell types, or analyze the ability of drug candidates to inhibit kinase activity. To do this, recombinant kinase substrates are immobilized on

membranes, incubated with purified kinase, and the substrates measured for the degree of phosphorylation.

Summary · ·

Meetings like this, packed with exciting discoveries and intriguing and interesting innovation, heavily emphasize the pace at which biotechnology is advancing, to the extent that the number of options for genomic and proteomic researchers can become overwhelming. Although data analysis is perhaps the greatest current concern for array users, an increasing challenge will be to determine the approaches and technology that really work, and to do it in a timely manner.

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Lal et al., 09/002,485, filed December 31, 1997 (PF-0459)

Exhibit "L" attached to Declaration of John C. Rockett, Ph.D.

A two-dimensional gel database of rat liver proteins useful in gene regulation and drug effects studies

A standard two-dimensional (2-D) protein map of Fischer 344 rat liver (F344MST3) is presented, with a tabular listing of more than 1200 protein species. Sodium dodecyl sulfate (SDS) molecular mass and isoelectric point have been established, based on positions of numerous internal standards. This map has been used to connect and compare hundreds of 2-D gels of rat liver samples from a variety of studies, and forms the nucleus of an expanding database describing rat liver proteins and their regulation by various drugs and toxic agents. An example of such a study, involving regulation of cholesterol synthesis by cholesterol-lowering drugs and a high-cholesterol diet, is presented. Since the map has been obtained with a widely used and highly reproducible 2-D gel system (the Iso-Dalt® system), it can be directly related to an expanding body of work in other laboratories.

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Abbreviations: CBB, Coomassie Brilliant Blue; CPK, creatine phosphokinase; 2-D, two-dimensional; 1EF, isoelectric focusing; MSN, master spot number; NP-40, Nonidet P-40, SDS, sodium dodecyl sulfate

1 Introduction

High-resolution two-dimensional electrophoresis of proteins, introduced in 1975 by O'Farrell and others [1-4], has been used over the ensuing 16 years to examine a wide variety of biological systems, the results appearing in more than 5000 published papers. With the advent of computerized systems for analyzing two-dimensional (2-D) gel images and constructing spot databases, it is also possible to plan and assemble integrated bodies of information describing the appearance and regulation of thousands of protein gene products [5, 6]. Creating such databases involves amassing and organizing quantitative data from thousands of 2-D gels, and requires a substantial commitment in technology and resources.

Given the long-term effort required to develop a protein database, the choice of a biological system takes on considerable importance. While in vitro systems are ideal for answering many experimental questions, especially in cancer research and genetics, our experience with cell cultures and tissue samples suggests that some in vivo approaches could have major advantages. In particular, we have noticed that liver tissue samples from rats and mice appear to show greater quantitative reproducibility (in terms of individual protein expression) than replicate cell cultures. This is perhaps a natural result of the homeostasis maintained in a complete animal vs. the well-known variability of cell cultures, the latter due principally to differences in reagents (e.g., fetal bovine serum), conditions (e.g., pH) and genetic "evolution" of cell lines while in culture. It is also more difficult to generate adequate amounts of protein from cell culture systems (particularly with attached cells), forcing the investigator to resort to radioisotope-based or silver-based staindetection methods. While these methods are more sensitive (sometimes much more sensitive) than the Coomassie Brilliant Blue (CBB) stain typically used for protein detection in "large" protein samples, they are generally more variable, more labor-intensive and, in the case of radiographic methods, may generate highly "noisy" images, due to the properties of the films used. By contrast, large protein samples can easily be prepared from liver using urea/Nonidet P-40 (NP-40) solubilization and stained with CBB, which has the advantage of being easily reproducible [8]. Finally, there remains the question of the "truthfulness" of many in vitro systems as compared to their in vivo analogs; how great are the changes caused by the introduction into a culture and the associated shift to strong selection for growth, and how do these affect experimental outcomes? Hence the apparent advantages of in vitro systems, in terms of experimental manipulation, may be counterbalanced by other factors relating to 2-D data quality.

There is a second important class of reasons for exploring the use of an in vivo biological system such as the liver. Historically, there have been two broad approaches to the mechanistic dissection of biochemical processes in intact cellular systems: genetics (a search for informative mutants) and the use of chemical agents (drugs and chemical toxins). Both approaches help us to understand complex systems by disrupting some specific functional element and showing us the result. With the development of techniques for genetic manipulation and cloning, the genetic approach can be effectively applied either in vitro or in vivo, although the in vitro route is usually quicker. The chemical approach can also be applied to either sort of biological system; here, however, the bulk of consistently acquired information is in experimental animals (rats and mice). While most biologists know a short list of compounds having specific, experimentally useful effects (e.g., inhibitors of protein synthesis, ionophores, polymerase inhibitors, channel blockers, nucleotide analogs, and compounds affecting polymerization of cytoskeletal proteins), there is a much larger number of interesting chemically-induced effects, most of them characterized by toxicologists and pharmacologists in rodent systems. Just as a thorough genetic analysis would involve saturating a genome with mutations, it is possible to imagine a saturating number of drugs, the analysis of whose actions would reveal the complete biochemistry of the cell. While organized drug discovery efforts usually target specific desired effects, the nature of the process, with its dependence on screening large numbers of compounds, necessarily produces many unanticipated effects. It is therefore reasonable to suppose that the required broad range of compounds necessary to achieve "biochemical saturation" may be forthcoming; in fact, it may already exist among the hundreds of thousands of compounds that failed to qualify as drugs.

Among organs, the liver is an obvious choice for the study of chemical effects because of its well-known plasticity and responsiveness. The brain appears to be quite plastic (e.g. [7]), but it is a complicated mixture of cell types requiring skillful dissection for most experiments. The kidney, while quite responsive, also presents a potentially confounding mixture of cell types. The liver, by contrast, is made up of one predominant cell type which is easy to solubilize: the hepatocyte, representing more than 95% of its mass. Most importantly, the liver performs many homeostatic functions that require rapid modulation of gene expression. It appears that most chemical agents tested affect gene expression in the liver at some dosage (N. Leigh Anderson, unpublished observations), an interesting contrast to our earlier work with lymphocytes, for example, which seem to be much less responsive. Such results conform to the expectation that cells with a homeostatic, physiological role should be more plastic than cells differentiated for a purpose dependent on the action of a limited number of specific genes.

The liver also allows the parallels between in vitro and in vivo systems to be examined in detail. Significant progress

has been made in the development of mouse, rat and human hepatocyte culture systems, as well as in precision-cut tissue slices. Using such an array of techniques, it is possible to assemble a matrix of mammalian systems including mouse and rat in vivo on one level and mouse, rat and human in vitro on a second level, and to compare effects between species and between systems. This approach allows us to draw informed conclusions regarding the biochemical "universality" of biological responses among the mammals, and to offer some insight into the validity of in vitro approaches for toxicological screening. We believe this data will be necessary if in vitro alternatives are to achieve wide usage in government-mandated safety testing of drugs, consumer products and industrial and agricultural chemicals.

A number of interesting studies have been published using 2-D mapping to examine effects in the rodent liver. A number of investigarors have made use of the technique to screen for existing genetic variants [8–11] or induced mutations [12–14], mainly in the mouse. This work builds on the wealth of genetic information available on the mouse and its established position as a mammalian mutation-detection system. While some studies of chemical effects have been undertaken in the mouse [15–17], most have used the rat [18–23]. The examination of the cytochrome p-450 system, in particular, has been carried out almost exclusively on the rat [24, 25].

These considerations lead us to conclude that rodent liver offers the best opportunity to systematically examine an array of gene regulation systems, and ultimately to build a predictive model of large-scale mammalian gene control. The basic underlying foundation of such a project is a reliable, reproducible master 2-D pattern of liver, to which ongoing experimental results can be referred. In this paper, we report such a master pattern for the acidic and neutral proteins of rat liver (pattern F344MST3). In future, this master will be supplemented by maps of basic proteins, and analogous maps of mouse and human liver.

2 Materials and methods

2.1 Sample preparation

Liver is an ideal sample material for most biochemical studies, including 2-D analysis. A sample is taken of approximately 0.5 g of tissue from the apical end of the left lobe of the liver. Solubilization is effected as rapidly as practical; a delay of 5-15 min appears to cause no major alteration in liver protein composition if the liver pieces are kept cold (e.g., on ice) in the interim. In the solubilization process, the liver sample is weighed, placed in a glass homogenizer (e.g., 15 mL Wheaton); 8 volumes of solubilizing solution*

The solubilizing solution is composed of 2% NP-40 (Sigma), 9 M urea (analytical grade, e.g., BDH or Bio-Rad), 0.5% dithiothreitol (DTT; Sigma) and 2% carrier ampholytes (pH 9-11 LKB: these come as a 20% stock solution, so 2% final concentration is achieved by making the final solution 10% 9-11 Ampholine by volume). A large batch of solubilizer (several hundred mL) is made and stored frozen at -80°C in aliquots sufficient to provide enough for one day's estimated sample preparation requirement. The solution is never allowed to become warmer than room temperature at any stage during preparation or thawing for use, since heating of concentrated urea solutions can produce contaminants that covalently modify proteins producing artifactual charge shifts. Once thawed, any unused solubilizer is discarded.

is added (i.e., 4 mL per 0.5 g tissue) and the mixture is homogenized using first the loose- and then then the tight-fitting glass pestle. This takes approximately 5 strokes with each pestle and is carried out at room temperature because urea would crystallize out in the cold. Once the liver sample is thoroughly homogenized in the solubilizer, it is assumed that all the proteins are denatured (by the chaotropic effect of the urea and NP-40 detergent) and the enzymes inactivated by the high pH (-9.5). Therefore these samples may be kept at room temperature until they can be centrifuged or frozen as a group (within several hours of preparation). The samples are centrifuged for 6×10^6 g min (e.g., 500 000 × g for 12 min using a Beckman TL-100 centrifuge). The centrifuge rotor is maintained at just below room temperature (e.g., 15-20°C), but not too cold, so as to prevent the precipitation of urea. The centrifuge of choice is a Beckman TL-100 because of the sample tube sizes available, but any ultracentrifuge accepting smallish tubes will suffice. When an appropriate centrifuge is not available near the site of sample preparation, samples can be frozen at -80°C and thawed prior to centrifugation and collection of supernatants. Each supernatant is carefully removed following centrifugation and aliquoted into at least 4 clean tubes for storage. This is done by transferring all the supernatant to one clean tube, mixing this gently (to assure homogeneous composition) and then dividing it into 4 aliquots. The aliquots are frozen immediately at -80°C. These multiple aliquots can provide insurance against a failed run or a freezer breakdown.

2.2 Two-dimensional electrophoresis

Sample proteins are resolved by 2-D electrophoresis using the 20 × 25 cm Iso-Dalt® 2-D gel system ([26-29]; produced by LSB and by Hoefer Scientific Instruments, San Francisco) operating with 20 gels per batch. All first-dimensional isoelectric focusing (IEF) gels are prepared using the same single standardized batch of carrier ampholytes (BDH 4-8A in the present case, selected by LSB's batchtesting program for rat and mouse database work**). A 10 μL sample of solubilized liver protein is applied to each gel, and the gels are run for 33 000 to 34500 volt-hours using a progressively increasing voltage protocol implemented by a programmable high-voltage power supply. An Angelique" computer-controlled gradient-casting system (produced by LSB) is used to prepare second-dimensional sodium dodecyl sulfate (SDS) polyacrylamide gradient slab gels in which the top 5% of the gel is 11%T acrylamide, and the lower 95% of the gel varies linearly from 11% to 18%T.

This system has recently been modified so as to employ a commercially available 30.8 %T acrylamide/N,N-methylenebisacrylamide prepared solution (thus avoiding the handling of the solid acrylamide monomer) and three additional stock solutions: buffer (made from Sigma pre-set Tris), persulfate and N,N,N',N'-tetramethylethylenediamine (TEMED). Each gel is identified by a computer-printed filter paper label polymerized into the lower left corner of the gel. First-dimensional IEF tube gels are loaded

directly (as extruded) onto the slab gels without equilibration, and held in place by polyester fabric wedges (Wedgies", produced by LSB) to avoid the use of hot agarose. Second-dimensional slab gels are run overnight, in groups of 20, in cooled DALT tanks (10°C) with buffer circulation. All run parameters, reagent source and lot information, and notations of deviation from expected results are entered by the technician responsible on a detailed, multi-page record of the experiment.

2.3 Staining

Following SDS-electrophoresis, slab gels are stained for protein using a colloidal Coomassie Blue G-250 procedure in covered plastic boxes, with 10 gels (totalling approximately 1 L of gel) per box. This procedure (based on the work of Neuhoff [30, 31]) involves fixation in 1.5 L of 50% ethanol and 2% phosphoric acid for 2h, three 30 min washes. each in 2L of cold tap water, and transfer to 1.5L of 34% methanol, 17% ammonium sulfate and 2% phosphoric acid for 1 h. followed by the addition of a gram of powdered Coomassie Blue G-250 stain. Staining requires approximately 4 days to reach equilibrium intensity, whereupon gels are transferred to cool tap water and their surfaces rinsed to remove any particulate stain prior to scanning. Gels may be kept for several months in water with added sodium azide. The water washes remove ethanol that would dissolve the stain (and render the system noncolloidal, with high backgrounds). The concentrated ammonium sulfate and methanol solution is diluted by equilibration with the water volume of the gels to automatically achieve the correct final concentrations for colloidal staining. Practical advantages of this staining approach can be summarized as follows: (i) the low, flat background makes computer evaluation of small spots (max OD < 0.02) possible, especially when using laser densitometry; (ii) up to 1500 spots can be reliably detected on many gels (e.g., rat liver) at loadings low enough to preserve excellent resolution; and (iii) reproducibility appears to be very good: at least several hundred spots have coefficients of reproducibility less than 15%. This value is at least as good as previous CBB methods, and significantly better than many silver stain systems.

2.4 Positional standardization

The carbamylated rabbit muscle creatine phosphokinase (CPK) standards [32] are purchased from Pharmacia and BDH. Amino acid compositions, and numbers of residues present in proteins used for internal standardization, are taken from the Protein Identification Resource (PIR) sequence database [33].

2.5 Computer analysis

Stained slab gels are digitized in red light at 134 micron resolution, using either a Molecular Dynamics laser scanner (with pixel sampling) or an Eikonix 78/99 CCD scanner. Raw digitized gel images are archived on high-density DAT tape (or equivalent storage media) and a greyscale videoprint prepared from the raw digital image as hard-copy backup of the gel image. Gels are processed using the Kepler® software system (produced by LSB), a commercially available workstation-based software package built on

^{**} This material (succeeding certified batches of which are available from Hoefer Scientific Instruments) has the most linear pH gradient produced by any ampholyte tested except for the Pharmacia wide range (which has an unacceptable tendency to bind high-molecular weight acidic proteins, causing them to streak).

some of the principles of the earlier TYCHO system [34–41]. Procedure PROC008 is used to yield a spotlist giving position, shape and density information for each detected spot. This procedure makes use of digital filtering, mathematical morphology techniques and digital masking to remove the background, and uses full 2-D least-squares optimization to refine the parameters of a 2-D Gaussian shape for each spot. Processing parameters and file locations are stored in a relational database, while various log files detailing operation of the automatic analysis software are archived with the reduced data. The computed resolution and level of Gaussian convergence of each gel are inspected and archived for quality control purposes.

Experiment packages are constructed using the Kepler experiment definition database to assemble groups of 2-D patterns corresponding to the experimental groups (e.g., treated and control animals). Each 2-D pattern is matched to the appropriate "master" 2-D pattern (pattern F344MST3 in the case of Fischer 344 rat liver), thereby providing linkage to the existing rodent protein 2-D databases. The software allows experiments containing hundreds of gels to be constructed and analyzed as a unit, with up to 100 gels displayed on the screen at one time for comparative purposes and multiple pages to accommodate experiments of > 1000 gels. For each treatment, proteins showing significant quantitative differences vs. appropriate controls are selected using group-wise statistical parameters (e.g., Student's t-test, Kepler® procedure STUDENT). Proteins satisfying various quantitative criteria (such as P <0.001 difference from appropriate controls) are represented as highlighted spots onscreen or on computer-plotted protein maps and stored as spot populations (i.e., logical vectors) in a liver protein database. Quantitative data (spot parameters, statistical or other computed values) are stored as real-valued vectors in the database. Analysis of coregulation is performed using a Pierson product-moment correlation (Kepler procedure CORREL) to determine whether groups of proteins are coordinately regulated by any of the treatments. Such groups can be presented graphically on a protein map, and reported together with the statistical criteria used to assess the level of coregulation. Multivariate statistical analysis (e.g., principal components' analysis) is performed on data exported to SAS (SAS Institute).

2.6 Graphical data output

Graphical results are prepared in GKS and translated within Kepler® into output for any of a variety of devices. Linedrawing output is typically prepared as Postscript and printed on an Apple Laserwriter. Detailed maps presented here have been generated using an ultra-high-resolution Postscript-compatible Linotronic output device. Greyscale graphics are reproduced from the workstation screen using a Seikosha videoprinter. Patterns are shown in the standard orientation, with high molecular mass at the top and acidic proteins to the left.

2.7 Experiment LSBC04

In the study described here 12-week-old Charles River male F344 rats were used. Diets were prepared at LSB, based on a Purina 5755M Basal Purified Diet. Lovastatin and cholestyramine were obtained as prescription pharma-

ceuticals, ground and mixed with the diet at concentrations of 0.075% and 1%, respectively. The high cholesterol diet was Purina 5801M-A (5% cholesterol plus 1% sodium cholate in the control diet). Animal work was carried out by Microbiological Associates (Bethesda, MD). Animals were acclimatized for one week on the control diet, fed test or control diets for one week, and sacrificed on day 8. Average daily doses of lovastatin and cholestyramine in appropriate groups were 37 mg/kg/day and 5 g/kg/day, respectively, based on the weight of the food consumed. Liver samples were collected and prepared for 2-D electrophoresis according to the standard liver protocol (homogenization in 8 volumes of 9 M urea, 2% NP-40, 0.5% dithiothreitol, 2% LKB pH 9-11 carrier ampholytes, followed by centrifugation for 30 min at $80\,000 \times g$). Kidney, brain and plasma samples were frozen. Gels were run as described above, and the data was analyzed using the Kepler system. Gels were scaled, to remove the effect of differences in protein loading, by setting the summed abundances of a large number of matched spots equal for each gel (linear scaling).

3 Results and discussion

3.1 The rat liver protein 2-D map

F344MST3 is a standard 2-D pattern of rat liver proteins, based on the Fischer 344 strain. This pattern was initiated from a single 2-D gel and extensively edited in an experiment comparing it to a range of protein loads, so as to include both small spots and well-resolved representations of high-abundance spots. More than 700 rat liver 2-D patterns have been matched to F344MST3 in a series of drug effects and protein characterization experiments, and numerous new spots (induced by specific drugs, for instance) have been added as a result. A modified version including additional spots present in the Sprague-Dawley outbred rat has also been developed (data not shown). Figure 1 shows a greyscale representation and Fig. 2 a schematic plot of the master pattern. More than 1200 spots are included, most of which are visible on typical gels loaded with 10 µL of solubilized liver protein prepared by the standard method and stained with colloidal Coomassie Blue. Master spot numbers (MSN's) have been assigned to all proteins, and appear in the following figures, each showing one quadrant of the pattern. Figure 3 shows the upper left (acidic, high molecular mass) quadrant, Fig. 4 the upper right (basic, high molecular mass) quadrant, Fig. 5 the lower left (acidic, low molecular mass) quadrant, and Fig. 6 the lower right (basic, low molecular mass) quadrant. The quadrants overlap as an aid to moving between them. The gel position (in 100 micron units), isoelectric point (relative to the CPK internal pI standards) and SDS molecular mass (from the calibration curve in Fig. 8) are listed for each spot (Table 1). Because of the precision of the CPK-p/values, these parameters can be used to relate spot locations between gel systems more reliably than using pl measurements expressed as pH. A major objective of current studies is the identification of all major spots corresponding to known liver proteins, as well as rigorous definitions of subcellular organelle contents. Of particular interest to us is the parallel development of identifications in the rat and mouse liver maps, allowing detailed comparisons of gene expression effects in the two systems. The results of these studies will be presented systematically in a later edition of this database, but we include here a useful series of 22 orienting identifications as an aid to other users of the rat liver pattern (Table 2).

3.2 Carbamylated charge standards, computed pls and molecular mass standardization

We have previously shown that the use of a system of closely-spaced internal pI markers (made by carbamylating a basic protein) offers an accurate and workable solution to the problem of assigning positions in the pI dimension [32]. The same system, based on 36 protein species made by carbamylating rabbit muscle CPK, has been used here to assign pI's to most rat liver acidic and neutral proteins. The standards were coelectrophoresed with total liver proteins, and the standard spots added to a special version of the master pattern F344MST3. The gel X-coordinates of all liver protein spots lying within the CPK charge train were then transformed into CPK pI positions by interpolation between the positions of immediately adjacent standards (Table 1) using a Kepler® vector procedure.

It has proven possible to compute fairly accurate pl values for many proteins from the amino acid composition [42]. We have attempted here to test a further elaboration of this approach, in which we computed pI's for the CPK standards themselves, based on our knowledge of the rabbit muscle CPK sequence and the fact that adjacent members of the charge train typically differ by blockage of one additional lysine residue (Table 3). We compared these values to similar computed pl's for an additional set of carbamylated standards made from human hemoglobin beta chains and a series of rat liver and human plasma proteins of known position and sequence (Fig. 7, Table 4). The result demonstrates good concordance between these systems. Two proteins show significant deviations: liver fatty-acid binding protein (FABP; #1 in Table 4) and protein disulphide isomerase (#20 in the table). The FABP spot present on F344MST3 may represent a charge-modified version of a more basic parent spot closer to the expected pl, not resolved in the IEF/SDS gel. Of particular importance is the fact that, by comparing computed prs of sequenced but unlocated proteins with the CPK pls, we can assign a probable gel location without making any assumptions regarding the actual gel pH gradient. This offers a useful shortcut, given the vagaries of pH measurement on small diameter IEF gels. We have used this approach to compute the CPK pr s of all rat and mouse proteins in the PIR sequence database, as an aid to protein identification (data not shown).

In order to standardize SDS molecular weight (SDS-MW), we have used a standard curve fitted to a series of identified proteins (Fig. 8). Rather than using molecular mass per se, we have elected to use the number of amino acids in the polypeptide chain, as perhaps a better indication of the length of the SDS-coated rod that is sieved by the second dimension slab. The resulting values were multiplied by 112 (the weighted average mass of amino acids in sequenced proteins) to give predicted molecular masses. Because we use gradient slabs, we have not constrained the fitted curve to conform to any predetermined model; rather we tried many equations and selected the best using the program "Tablecurve" on a PC. The equation chosen was y = a + bx + c/x, where y is the number of residues, x is the gel

Y coordinate, a is 511.83, b is -0.2731 and c is 33183801. The resulting fit appears to be fairly good over a broad range of molecular mass.

3.3 An example of rat liver gene regulation: Cholesterol metabolism

Experiment LSBC04 was designed as a small-scale test of the regulation of cholesterol metabolism in vivo by three agents included in the diet: lovastatin (Mevacor®, an inhibitor of HMG-CoA reductase); cholestyramine (a bile acid sequestrant that has the effect of removing cholesterol from the gut-liver recirculation); and cholesterol itself. The first two agents should lower available cholesterol and the third should raise it, allowing manipulation of relevant gene expression control systems in both directions. Such an experiment offers an interesting test of the 2-D mapping system since most of the pathway enzymes are present in low abundance, many are membrane-bound and difficult to solubilize, and the pathway itself is complex. Approximately 1000 proteins were separated and detected in liver homogenates. Twenty-one proteins were found to be affected by at least one treatment, and these could be divided into several coregulated groups.

3.3.1 MSN 413 (putative cytosolic HMG-CoA synthase) and sets of spots regulated coordinately or inversely

One group of spots (including a spot assigned to the cytosolic HMG-CoA synthase, MSN 413) showed the expected increase in abundance with lovastatin or cholestyramine, the synergistic further increase with lovastatin and cholestyramine, and a dramatic decrease with the high cholesterol diet. Spot number 413 is the most strongly regulated protein in the present experiment, showing a 5- to 10-fold induction after a 1 week treatment with 0.075 % lovastatin and 1% cholestyramine in the diet (Figs. 9 and 10). Its expression follows precisely the expectation for an enzyme whose abundance is controlled by the cholesterol level; it is progressively increased from the control levels by cholestyramine, lovastatin and lovastatin plus cholestyramine, and it sinks below the threshold of detection in animals fed the high cholesterol diet. This spot has been tentatively identified as the cytosolic HMG-CoA synthase, based on a reaction with an antiserum to that protein provided by Dr. Michael Greenspan at Merck Sharp & Dohme Research Laboratories. This enzyme lies immediately before HMG-CoA reductase in the liver cholesterol biosynthesis pathway, and is known to be co-regulated with it. Spot 413 has an SDS molecular weight of about 54 000 and a CPK pl of -11.4, in reasonably close agreement with a molecular weight of 57300 and a CPK pl of -15.7 computed from the known sequence of the hamster enzyme [43].

Using a classical product-moment correlation test (Kepler procedure CORREL), a series of five additional spots was found to be coregulated with 413. The level of correlation was exceedingly high (> 95%). Two of these, 1250 and 933, are at similar molecular weights and approximately one charge more acidic than 413 (Fig. 9), indicating that they may be covalently modified forms of the 413 polypeptide. This suspicion is strengthened by the observation that both spots are also stained by the antibody to cytosolic HMG-CoA synthase. The remaining three correlated spots appear

to comprise an additional related pair (1253 and 1001) of around 40 kDa and a single spot (1119) of around 28 kDa. Because these two presumed proteins are present at substantially lower abundances than 413, and because the cytosolic HMG-CoA synthase is reported to consist of only one type of polypeptide, they are likely to represent other, very tightly coregulated enzymes. A second group of six spots was selected based on a regulatory pattern close to the inverse of that for spot 413 (MSN's 34, 79, 178, 182, 204, 347; data not shown). For these proteins, the lowest level of expression occurs with exposure to lovastatin plus cholestyramine and the highest level upon exposure to the high-cholesterol diet. Spots 182 and 79 are highly correlated and lie about one charge apart at the same molecular weight; they may thus be isoforms of a single protein. The other four spots probably represent additional enzymes or subunits.

3.3.2 MSN 235 and coregulated spots

A third group of five spots, mainly comprised of mitochondrial proteins including putative mitochondrial HMG-CoA synthase spots, showed a modest induction by lovastatin alone, but little or no effect with any of the other treatments (including the combination of lovastatin and cholestyramine; Fig. 12). This result is intriguing because lovastatin was expected to affect only the regulation of enzymes of cholesterol synthesis, which is entirely extra-mitochondrial. Three of the spots (235, 134, 144) form a closelypacked triad at approximately 30 kDa, and are likely to represent isoforms of one protein. All three spots are stained by an antibody to the mitochondrial form of HMG-CoA synthase obtained from Dr. Greenspan. Subcellular fractionation indicates a mitochondrial location. The other two spots (633 at about 38 kDa and 724 at about 69 kDa) are each present at lower abundance than the members of the triad.

3.3.3 An example of an anti-synergistic effect

A sixth spot (367) shows strong induction by lovastatin (two- to threefold), and about half as much induction with lovastatin plus cholestyramine, but without sharing the animal-animal heterogeneity pattern of the 235-set (Fig. 13). This protein is also mitochondrial, and represents the clearest example of an anti-synergistic effect of lovastatin and cholestyramine. The existence of such an effect demonstrates that lovastatin and cholestyramine do not act exclusively through the same regulatory pathway.

3.3.4 Complexity of the cholesterol synthesis pathway

Taken together, these results suggest that treatment with lovastatin alone can affect both cytosolic and mitochondrial pathways using HMG-CoA, while cholestyramine, on the other hand, either alone or in combination with lovastatin, produces a strong effect on the putative cytosolic pathway, but little or no effect on the putative mitochondrial pathway. An explanation for this difference may lie in lovastatin's effect on levels of HMG-CoA and related precursor compounds that are exchanged between the cytosol and the mitochondrion, whereas cholestyramine should affect only the cytosolic pathways directly controlled by cholesterol and bile acid levels. It remains to be explained why some

proteins of the putative mitochondrial pathway are so much more variable in their expression in all groups. An examination of all the coregulated groups suggests that quantitative statistical techniques can extract a wealth of interesting information from large sets of reproducible gels. The abundance of spots in the 413 coregulation group, for example, shows an amazing level of concordance in their relative expression among the five individuals of the lovastatin and cholestyramine treatment group. This effect is not due to differences in total protein loading, since they have already been removed by scaling, and since proteins with quite different regulation patterns can be demonstrated (e.g., Fig. 13). Such effects raise the possibility that many gene coregulation sets may be revealed through the study of a sufficiently large population of control animals (i.e., without any experimental manipulation). This approach, exploiting natural biological variation in protein expression instead of drug effects, offers an important incentive for the construction of a large library of control animal patterns.

4 Conclusions

Because of the widespread use of rat liver in both basic biochemistry and in toxicology, there is a long-term need for a comprehensive database of liver proteins. The rat liver master pattern presented here has proven to be an accurate representation of this system, having been matched to more than 700 gels to date. As the number of proteins identified and the number of compounds tested for gene expression effects grows, we expect this database to contribute valuable insights into gene regulation. Its practical utility in several areas of mechanistic toxicology is already being demonstrated.

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5 References

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6 Addendum 1: Figures 1-13

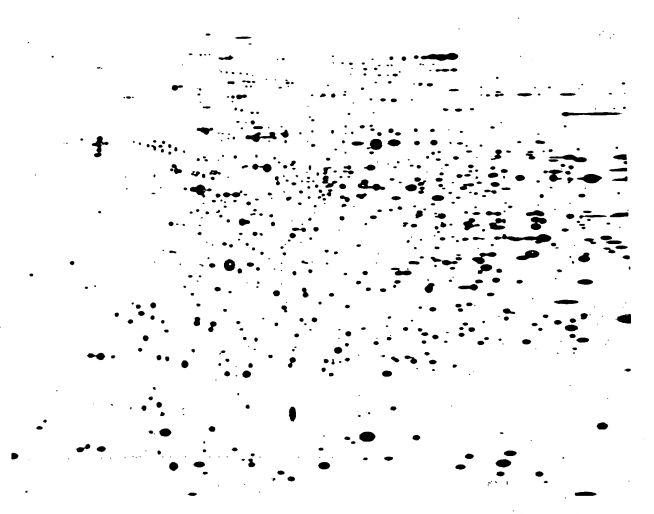


Figure 1. Synthetic representation of the standard rat liver 2-D master pattern, rendered as a greyscale image using a videoprinter.

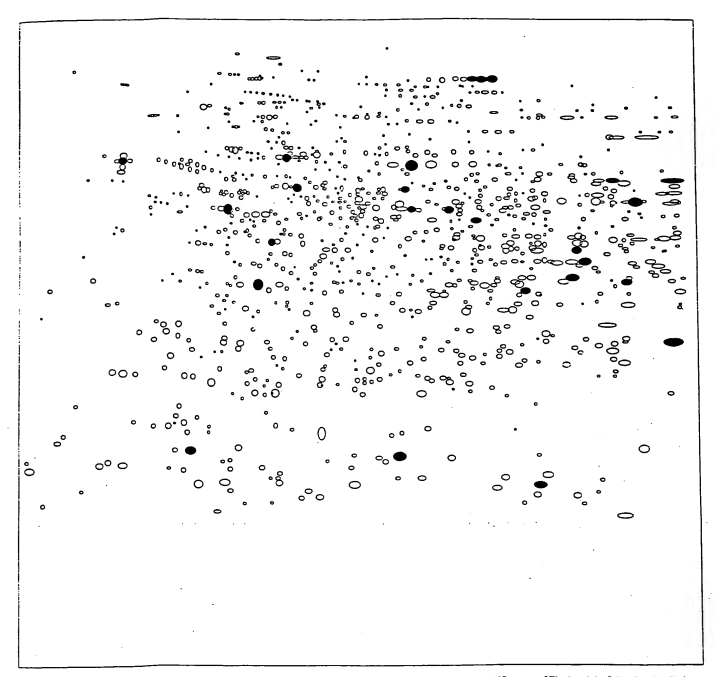


Figure 2. Schematic representation of the master pattern (the same as Fig. 1), useful as an aid in relating specific areas of Fig. 1 and the following detailed quadrants.

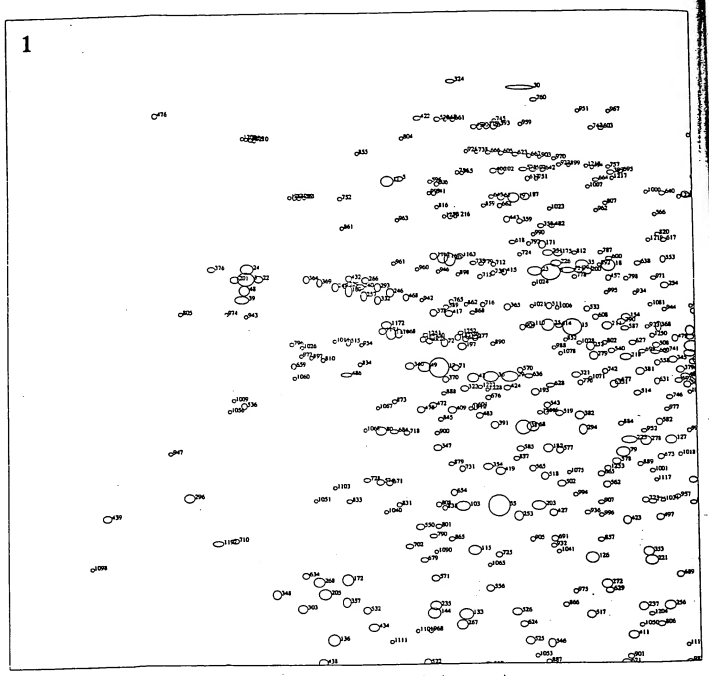


Figure 3. Upper left (high molecular weight, acidic) quadrant (#1) of the rat liver map, showing spot numbers.

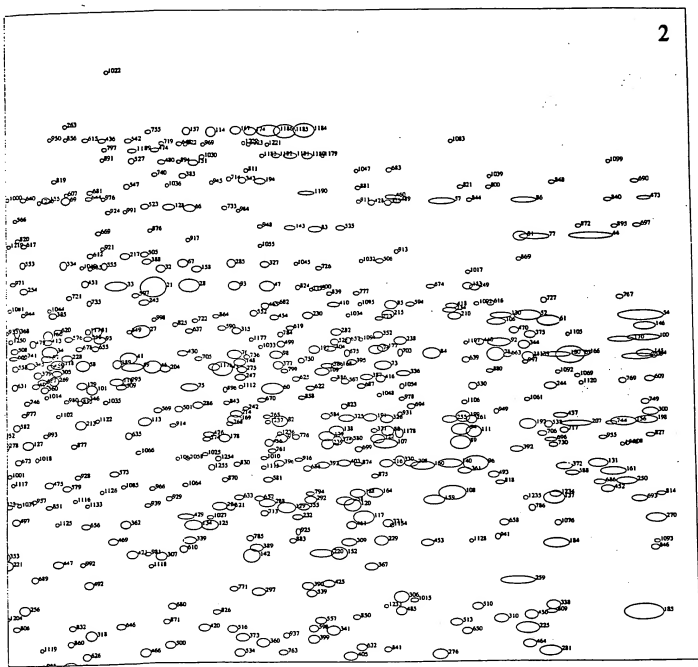


Figure 4. Upper right (high molecular weight, basic) quadrant (#2) of the rat liver map, showing spot numbers.

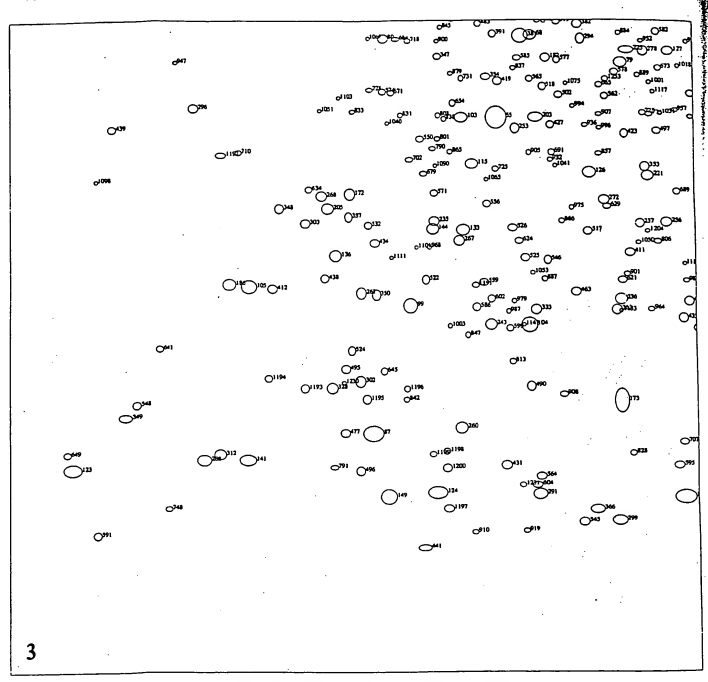


Figure 5. Lower left (low molecular weight, acidic) quadrant (#3) of the rat liver map, showing spot numbers.

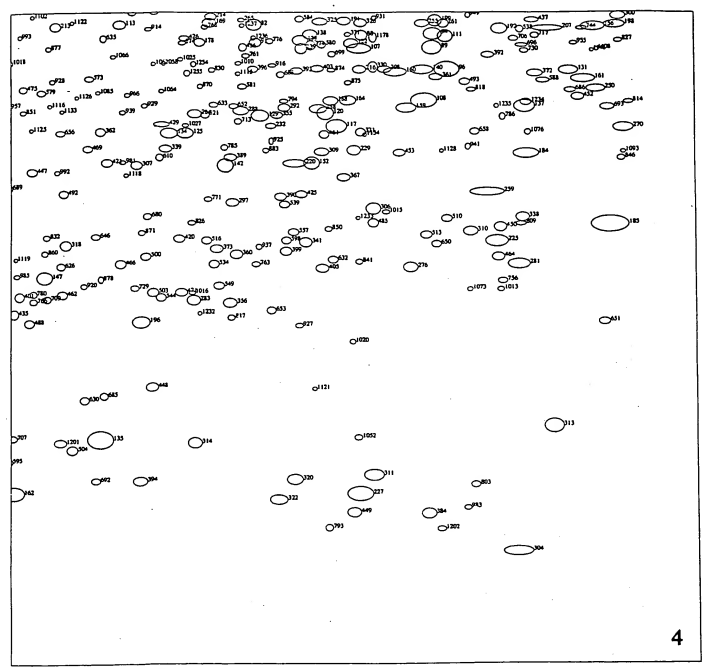
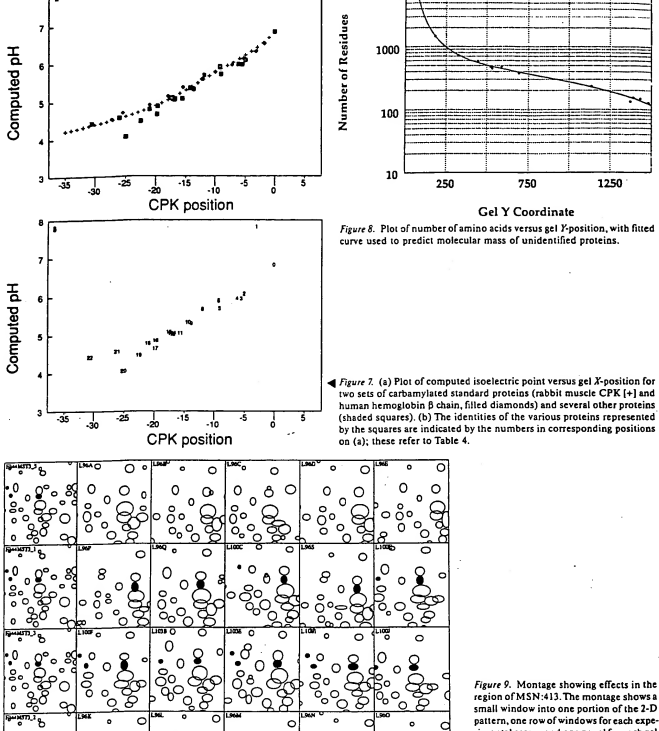


Figure 6. Lower right (low molecular weight, basic) quadrant (#4) of the rat liver map, showing spot numbers.



O

10000

region of MSN:413. The montage shows a small window into one portion of the 2-D pattern, one row of windows for each experimental group, and one panel for each gel in the experiment. The left-most pattern in each row is a group-specific copy of the master pattern followed by the patterns for the five individual rats in the group. The highlighted protein spots (filled circles) are spot 413 (on the right of each panel; identified as cytosolic HMG-CoA synthase) and two modified forms of it (1250 and 933). From the top, the rows (experimental groups) are: high cholesterol, controls, cholestyramine, lovastatin, and lovastatin plus cholestyramine.

Regulation of Rat Liver 413

(Putative Cytosolic HMG-CoA Synthase, 53kd)
Test Compounds in Diet

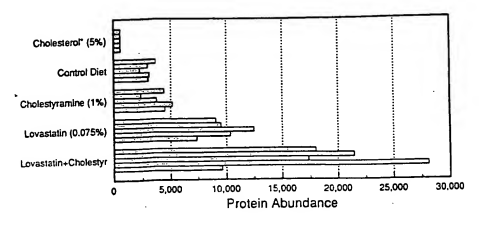


Figure 10. Bargraph showing the quantitative effects of various treatments on the abundance of MSN:413 (cytosolic HMG-CoA synthase) in the gels of Fig. 9.

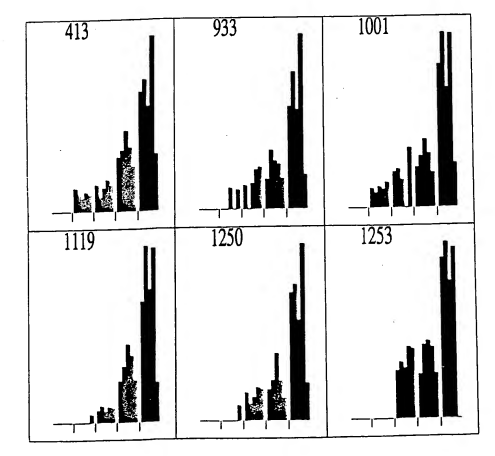


Figure 11. Bargraphs of a series of six coregulated spots including MSN:413. In the bargraphs, the abundances of the appropriate spot (master spot number shown at the top of the panel) in each animal are shown. The five five-animal groups are in the order (left to right): high cholesterol, controls, cholestyramine, lovastatin, and lovastatin plus cholestyramine. Each bar within a group represents one experimental animal liver (one 2-D gel). Note the correlated expression of the 6 spots, especially in the two far right (most strongly induced) groups.

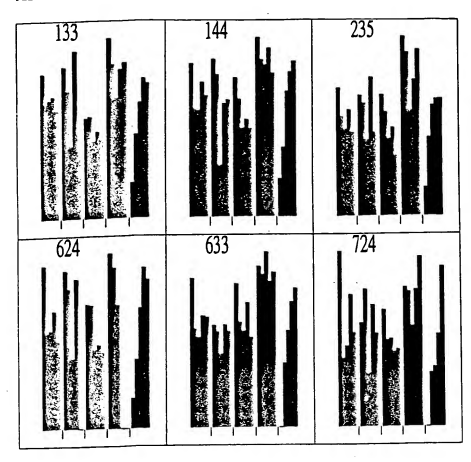


Figure 12. Data on a second coregulated group of spots, presented as in Fig. 11. The fourth experimental group (lovastatin) shows a modest induction, while the fifth group (lovastatin plus cholestyramine) does not.

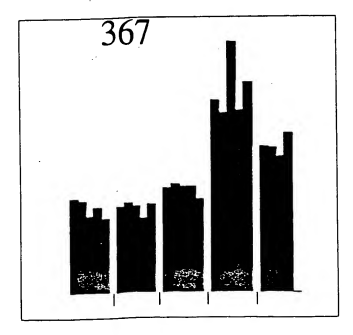


Figure 13. Data on spot MSN:367, presented as in Fig. 11. This protein shows unambiguously the anti-synergistic effect of lovastatin and cholestyramine (fifth group) as compared to lovastatin (fourth group). This response contrasts strongly with the regulation pattern seen in Fig. 11.

				in the rat liver	-				20212:	1.00			00:4:	656:5:
ISN	X	<u> </u>	CPKol	SDSMW	MSN	X	Y	CPKol	SDSMW	MSN	х	<u> </u>	CPKol	SDSMV
3	311	434	<-35.0	63,800	95	1119	536	-9.9	53,800	174	1364	183	-6.7	162,90
5	568	263	-24.3	102,900	96	1731	756	-2.0	40,700 51,600	175 177	825 1582	393 553	-15.7 -3.6	69,30 52,60
В	812 549	426 268	-16.0 -25.2	64,800 101,000	97 98	1033 1406	566 565	-11.4 -6.1	51,700	178	1321	710	-7.2	43,00
11 15	845	520	-15.3	55,200	99	578	1149	-23.8	25,000	179	1089	615	-10.4	48,30
17	629	589	-21.6	50,000	100	2004	538	>0.0	53,700	180	1866	567	-0.5	51,60
18	906	414	-14.0	66,300	101	1106	623	-10.1	47,900	181	411	295	-32.1	91,20
19	755	298	-17.5	90,200	102	482	455	-28.5	61,300 27,200	182 184	804 1860	730 896	-16.2 -0.6	42,00 34,50
20	649	403 448	-20.9 -8.7	67,900 62,100	103 104	665 773	830 1182	-20.2 -17.0	37,300 · 23,800	185	1997	1017	>0.0 >0.0	29,80
21 22	1204 332	434	<-35.0	63,800	105	312	1117	<-35.0	26,100	186	279	1113	<-35.0	26,30
23	787	424	-16.6	65,000	106	1769	509	-1.5	56,100	187	773	296	-17.0	90,80
24	313	417	<-35.0	66,000	107	1585	720	-3.6	42,500	188	1538	807	-4.2	38,40
25	807	516	-16.1	55,500 54,000	108 109	1692 1482	807 593	-2.4	38,300 49,700	191 192	1560 1818	674 687	-3.9 -0.9	44,90 44,20
27 28	1184 1263	524 446	-9.0 -8.0	54,900 62,400	110	778	516	-4.8 -16.9	55,500	193	1469	555	-5.0	52,40
29 29	743	605	-17.B	49,000	111	1728	700	-2.0	43,500	194	1380	266	-6.4	101,60
30	768	112	-17.2	348,600	113	1191	680	-8.9	44,500	195	784	632	-16.7	47,30
32	1216	417	-8.6	66,000	114	1298	185	-7.5	160,800	196	1227	1185	-8.4	23,70
33	1145	445 EEE	-9.5 -11.3	62,500 52,400	115 116	682 1146	907 610	-19.6 -9.5	34,100 48,700	197 198	667 2006	553 681	-20.1 >0.0	52,60 44,50
34 35	1037 863	555 412	-11.3 -14.9	66,600	117	1548	849	-9.5 -4.1	36,500	199	1711	674	-2.2	44,90
35 36	712	606	-18.7	48,900	118	1050	577	-11.1	50,800	200	872	424	-14.7	65,0
38	763	694	-17.3	43,800	120	1530	828	-4.3	37,400	201	292	435	<-35.0	63,7
39	304	470	<-35.0	59,800 51,400	121 122	638 1572	423 712	-15.4 -3.8	65,200 42,900	202 203	736 786	253 829	-18.0 -16.7	107,84 37,44
41 42	1165 684	569 607	-9.2 -19.6	48,800	123	23	1433	<-35.0	15,300	204	1224	589	-8.5	50,0
43	1318	589	-7.3	50,000	124	621	1474	-21.9	13,900	205	439	983	-30.9	31,10
44	1924	362	-0.1	74,600	125	1298	862	-7.5	36,000	206	1994	571	>0.0	51,3
46	1203	586	-8.7	50,200	126	872	921	-14.7	33,500 42,600	207	1895 240	687 1418	-0.3 <-35.0	44,20 15,80
47 48	1391 309	447 454	-6.3 <-35.0	62,300 61,500	127 128	1000 1229	717 311	-12.0 -8.4	86,100	210	1700	499	-2.3	57,0
40 49	605	587	-22.5	50,100	129	1422	832	-5.8	37,30C	211	902	517	-14.1	55,4
50	621	535	-21.8	53,900	130	1776	499	-1.4	57,000	213	1087	684	-10.4	44.4
51	1113	522	-10.0	55,000	131	1930	757	-0.1	40,700	214	1340	668	-7.0	45,2
52	1820	499 177	-0.9 -18.3	57,000 170,800	132 133	660 666	537 1019	-20.4 -20.2	53,800 29,700	215 216	1591 1585	495 755	-3.5 -3.6	57,3 40,7
53 54	725 2001	500	>0.0	56,900	134	1271	862	-7.9	36,000	217	1159	393	- 9 .3	69,3
55	722	830	-18.4	37,300	135	1161	1389	-9.3	16,800	218	931	572	-13.5	51,2
56	678	533	-19.8	54,100	136	453	1063	-29.7	28,100	219	713	177 911	-18.7	170,5
57	1682	302 580	-2.5 -10.3	89, 000 50,600	137 138	1858 1504	823 697	-0.6 -4.6	37,700 43,700	220 221	1479 965	927	-4.9 -12.8	33,9 33,3
58 59	1091 1171	585	-9.2	50,300	139	1488	707	-4.8	43,200	223	934	716	-13.5	42,7
60	1400	624	-6.2	47,800	140	1689	756	-2.4	40,700	225	1812	1045	-1.0	28,8
61	1853	508	-0.6	56,200	141	311	1417	<-35.0	15,800	226	821	411	-15.8	66,8
62	1688	567	-0.4	51,500 90,500	142 143	1366 1429	915 346	-6.7 -5.7	33,800 77,900	227 228	1586 1065	1483 567	-3.6 -10.8	13,6 51,6
65 66	735 1263	297 312	-18.1 -8.0	85, 900	144	615	1017	-22.1	29,800	229	1577	890	-3.7	34,8
67	1252	407	-8.1	67,300	145	2006	566	>0.0	51,600	230	1458	496	-5.2	57,3
68	779	692	-16.8	43,900	146	2006	518	>0.0	55,300	232	1440	849	-5.5	36,5
69	1064	296	-10.8	90,800	147	1070	1108	-10.7	26,500 50,800	234 235	1692 618	489 1004	-2.4 -22.0	57,9 30,3
71 72	656 638	589 545	-20.6 -21.2	50,000 53,100	148 149	1347 541	578 1481	-6.9 -25.7	13,700	236	920	1138	-13.7	30,3 25,4
72 73	1582	583	-3.6	50,400	150	1645	760	-2.8	40,500	237	952	1008	-13.1	30,2
74	1570	556	-3.8	52,300	151	1269	236	-7.9	117,000	238	1611	541	-3.2	53,5
75	1264	621	-8.0	48,000	152	1507	911	-4.5 -2.1	33,900 62,100	239 240	1489 501	720 448	-4.8 -27.7	42,5 62,1
76 77	1338	564 363	-7.0 -0.8	51,800 74,400	153 154	1722 932	448 503	-2.1 -13.5	62,100 56,600	240	1820	569	-27.7 -0.9	51,4
77 78	1833 1767	565	-0.6 -1.5	51,700	155	1031	294	-11.4	91,400	242	1357	658	-6.8	45,8
79	925	738	-13.6	41,600	156	1970	684	>0.0	44,400	243	711	1182	-18.7	23,8
80	534	698	-26.1	43,600	157	1258	183	-8.1	162,400	244	1855	621	-0.6 -8.0	48,0
81	1811	363	-1.0 -6.0	74,500 44,500	158 159	1275 1663	417 820	-7.8 -2.6	65,900 37,800	245 246		474 459	-8.9 -25.1	59,3 61,0
82 83	1412 1471	681 347	-5.0 -5.0	77,500	160	1034	527	-11.4	54,600	247	1348	604	-6.9	49,1
84	1662	563	-2.7	51,800	161	1953	771	>0.0	40,000	248	460	448	-29.3	62,
85	1596	479	-3.4	58,900	162	1020	1482	-11.6	13,700	249	1733	451	-1.9	61,8
86	1817	301	-0.9	89,100	164	1566	806	-3.8	38,400	250	1974 808	788 392		39,2 69,5
87	516	1371	-27.0 -3.5	17,400 43,600	166 167	1905 1340	565 181	-0.2 -7.0	51,700 164,900	251 252		553		52,
88 89	1589 1706	698 719	-3.5 -2.2	42,500	168	1506	583	-7.0 -4.6	50,400	253		848		36,
90	651	329	-20.8	81,700	169	1338	678	, -7.0	44,700	254	995	450	-12.1	61,9
91	1415	710	-6.0	43,000	170	1969	541	>0.0	53,500	255				44,0
92	1773	545	-1.4	53,200 62,300	171 172	800 476	378 958	-16.3 -28.7	71,800 32,100	256 257		1006 464		30,2 60,4
93	1338	446	-7.0											

Master table of proteins in the rat liver database, showing spot master number, gel position (x and y), isoelectric point relative to CPK standards, and predicted molecular mass (from the standard curve of Fig. 8).

The state of the s

MSN	х	Υ	CPKol	SDSMW	MSN	X	Υ	CPKøl	SDSMW	MSN	x	Y	CPKel	SDSMW
					245	1006	578	-11.9	50,800	426	1296	704	-7. 6	43,300
259	1796	961	-1.1	31,900 17,700	345 346	1006 1095	640	-10.3	46,800	427	810	843	-16.0	36,800
260	661	1361 679	-20.4 -2.0	44,600	347	625	728	-21.7	42,000	428	1565	303	-3.9	88,700
261 262	1725 496	1127	-28.0	25,800	348	361	983	-35.3	31,100	429	1259	847	-8.0 -8.1	36,600
263	1063	172	-10.9	177,400	349	110	1343	<-35.0	18,300	430 431	1253 734	562 1426	-0.1 -18.1	51, 900 15,500
265	1390	673	-6.3	45,000	350	521	1130	-26.7	25,700 48,100	432	483	433	-28.5	63,900
266	510	437	-27.3	63,400	351 352	912 1574	619 530	-13.9 -3.7	54,300	434	518	1041	-26.9	28,900
267	660	1038	-20.4	29,000 31,900	352	961	912	-12.9	33,900	435	1020	1170	-11.6	24,300
268	430	961 606	-31.0 -11.2	48,900	354	706	762	-18.9	40,400	436	1122	196	-9.8	147,600
269	1044 2019	853	>0.0	36,300	355	1450	830	-5.3	37,300	437	1870	673	-0.5	45,000
270 271	857	422	-15.0	65,200	356	1374	1152	-6.5	24,900	438	435 86	1102 847	-31.0 <-35.0	26,700 36,600
272	895	968	-14.2	31,700	357	474	997	-28.7	30,6 00 77,8 00	439 440	1740	544	-1.8	53,200
274	1292	712	-7.6	42,900	358 359	798 764	346 338	-16.3 -17.3	79,400	441	599	1571	-22.8	10,800
275	1350	590	-6.9 -2.6	49,900 27,100	360	1384	1068	-6.4	27,900	443	743	335	-17.8	80,100
276	1670	1089 538	-19.4	53,700	361	1713	769	-2.1	40,100	446	801	668	-16.2	45,200
277 278	688 961	718	-13.0	42,600	362	1161	859	-9.3	36,100	447	1050	926	-11.1	33,300
279	879	570	-14.5	51,300	363	914	1156	-13.8	24,800	448	1245 1576	1298 1516	-8.2 -3.7	19, 800 12,600
281	1848	1084	-0.7	27,300	364	412	435	-32.0	63,700 58,200	449 450	1818	1021	-0.9	29,600
282	1505	525	4.6	54,800 35,100	365 366	741 878	486 1503	-17.9 -14.6	13,000	451	1094	440	-10.3	63,100
283	1313	1147	-7.3 -7.3	25,100 37,400	367	1560	935	-3.9	33,000	452	1945	802	>0.0	38,600
284	1314 1332	829 408	-7.1	67,200	368	983	520	-12.4	55,200	453	1652	894	-2.8	34,600
285 286	1277	652	-7.8	46,100	369	434	441	-31.0	63,000	454	1403	500	-6.1	56,900 42,600
288	1391	824	-6.3	37,600	370	639	610	-21.2	48,700	456 457	1394 905	718 436	-6.3 -14.0	63,500
289	1147	579	-9.5	50,700	371	1587	860 762	-3.6 -0.5	36,100 40,400	459	1038	581	-11.3	50,500
290	925	511	-13.6	55,900 13,900	372 373	. 1875 1351	1059	-6.8	28,300	460	1598	294	-3.4	91,400
291	787	1476 818	-16.6 -5.1	37,800	374	1506	715	-4.6	42,700	461	1528	863	-4.3	35,900
292 293	1462 531	449	-26.3	62,000	375	1823	532	-0.9	54,200	462	1098	1137	-10.2	25,400
294	860	698	-14.9	43,600	376	254	417	<-35.0	65,900	463	849	1125 1072	-15.2 -0.9	25,800 27,800
295		609	-9.3	48,700	377	1409	583	-6.1	50,400 57,500	464 465	1814 1388	481	-6.3	58,700
296		814	<-35.0	38,000	378 379	621 1017	494 595	-21.8 -11.7	49,600	466	1194	1084	-8.9	27,300
297		979	-6.5 -13.9	31,300 12,400	381	953	598	-13.1	49,400	468	577	467	-23.9	60,100
299		1523 667	>0.0	45,300	382	856	674	-15.0	44,900	469	1140	888	-9.6	34,900
300 301	702	178	-19.0	169,200	383	1252	258	-8.1	105,300	470	1797	524	-1.1	54,800 25,500
302		1280	-28.1	20,400	384	1699	1518	-2.3	12,500	471 472	1293 618	1133 655	-7.6 -21.9	46,000
303		1008	-32.6	30,100	385	1042 1490	493 583	-11.2 -4.7	57,500 50,400	473	2009	299	>0.0	89,900
304		1585	-0.7 -11.1	10,300 49,800	386 387	1554	603	-4.0	49,100	474	1205	215	-8.7	131,300
305		593 989	-3.3	30,900	388	1193	404	-8.9	67,700	475	1035	788	-11.4	39,200
306 307		916	-8.5	33,700	389	1374	902	-6.5	34,300	476	160	155	<-35.0	207,600
308		755	-3.0	40,700	390	1456	969	-5.2	31,700	477 478	469 599	1370 662	-28.9 -22.8	17,400 45,600
309	1524	892	-4.4	34,700	391	718	690	-18.5	44,000 41,900	479	1009	540	-11.8	53,500
310		1028	-1.5	29,400 14,700	392 393	1799 1482	732 758	-1.1 -4.8	40,600	480		235	-8.6	117,400
311		1451 1408	-3.3 <-35.0	16,100	394	1227	1461	-8.4	14,400	482	816	346	-15.9	77,800
312 313		1365	-0.3	17,600	395	1530	577	-4.3	50,800	483		673	-19.3	44,900
314		1395		16,600	396	1410	755		40,800	485		1013 599	-3.3 -28.6	30,000 49,300
315		523	-7.0	54,900	397	912	256		106,400 28,100	486 487		607		48,800
→ 318		1053		28,500	399 400		1063 450		61,900	488		1186		23,700
320		1459		14,400 49,1 0 0	401	1029	1140		25,300	489		301		89,200
32 ⁻ 32 ⁻		603 1494		13,300	403		754		40,800	490		1289		20,100
32		626		47,700	404		554		52,500	491		178		169,300
32		101		420,500	405		1092		27,100	492		964 776		31,800 39,700
32		675		44,800	406		252		108,000	493 494				110,700
32		677		44,700	409		663		45,500 59,000	495				21,200
32		409		67,000 20,100	410 411		478 1057		28,300	496		1436		15,200
32		1291 751		40,900	412		1120		26,000	497	7 980			36,400
33		697		43,700	413		538		53,700	499				53,100
33 ⁻ 33 ⁻		471		59,600	415	737	425		64,900	500				27,800 45,700
33		1156	-16.7	24,700	416				48,900 57,300	50° 50°				
33	1059			67,300	417				57,300 58,600	50. 50.				
33				88,500 49,400	418 419				40,000	50-				
33				49,400 30,300	420		1041		28,900	50				69,700
33		1004 888		34,900	421		912		33,900	50				
33' 34		585		50,300	422			-22.8	193,700	50				
34		_		28,700	423	929				50				
34		265	-6.8	102,200	424					50 51				
34		549	-0.9	52.800	425	1490	965	5 -4.7	31.800	31	0 1730	, 100	-2.0	50,250

MSN	Х	Y	CPKol	SDSMW	MSN	X	Y	CPKpl	SDSMW	MSN	X	Y	CPKol	SDSMW
F11	809	484	-16.0	58,400	596	619	269	-21.9	100,500	674	1661	448	-27	62,100
511 512	1099	533	-10.2	54,100	597	1176	461	-9.1	60,700	675	1523	562	-4.4	51,900
513	1696	1034	-2.3	29,200	598	1465	1044	-5.0	28,800	676	708	642	-18.8	46,700
514	948	636	-13.2	47,100	599	741	1188	-17.9	23, 600 68,000	677 678	919 1085	615 551	-13.7 -10.5	48,300 52,700
515	481 1334	543 1044	-28.5 -7.1	53, 400 28,800	600 601	907 687	402 658	-14.0 -19.5	45,800	679	600	923	-22.7	33,400
516 517	868	1021	-14.8	29,700	602	712	1138	-18.7	25,400	680	1237	1004	-8.3	30,300
518	798	779	-16.3	39,600	603	898	181	-14.1	165,200	681	1103	283	-10.1	95,100
519	822	670	-15.7	45,100	604	783	1461	-16.7	14,400	682	1406	477	-6.1 -3.4	59,100
520	632	165 830	-21.5 -7.1	189,000 37, 30 0	605 606	736 629	223 273	-18.0 -21.6	125, 300 98,7 00	683 684	1596 555	249 699	-24.8	109,800 43,500
521 522	1332 603	1104	-22.6	26,600	607	1064	286	-10.8	94,000	685	1167	1313	-9.2	19,300
523	1190	309	-8.9	86,800	608	883	503	-14.5	56,700	686	1932	790	0.0	39,100
524	479	1226	-28.6	22,300	609	2012	610	>0.0	48,700	687 688	1545 1456	619 764	-4.1 -5.2	48,100 40,300
525	768 747	1066 1016	-17.2 -17.7	28, 000 29, 800	610 612	1255 1103	903 391	-8.1 -10.1	34,200 69,600	689	1011	953	-5.2 -11.8	32,300
526 527	1170	231	-9.2	119,600	613	778	265	-16.9	102,000	690	1995	270	>0.0	100,200
528	1502	542	-4.6	53,400	614	.824	518	-15.7	55,400	691	812	888	-16.0	34,900
530	1728	620	-2.0	48,000	615	1095	195	-10.3	149,100	692	1154	1461	-9.4	14,400
532	507	1011	-27.4 -14.7	30, 000 57, 900	61 6 617	1759 994	478 372	-1.6 -12.1	59, 000 72, 900	693 694	1993 1628	819 656	>0.0 -3.0	37,800 45,900
533 534	870 1347	489 1085	-6.9	27,300	618	751	374	-17.6	72,400	695	928	254	-13.6	107,000
535	1513	346	-4.5	77,800	619	1429	518	-5.7	55,300	696	1854	715	-0.6	42,700
536	308	654	<-35.0	46,000	620	1050	520	-11.1	55,200	697	1997	345	>0.0	78,000
538	1851	689 982	-0.7 -5.1	44,100 31,100	621 622	923 1462	1105 622	-13.7 -5.1	26,600 47,900	698 699	957 1540	563 730	-13.0 -4.2	51,800 42,000
539 540	1463 909	561	-13.9	52,000	623	759	225	-17.4	124,000	702	577	900	-23.8	34,400
541	625	289	-21.7	93,100	624	758	1038	-17.4	29,000	703	1610	562	-3.2	51,900
542	1164	198	-9.2	146,200	625	1438	606	-5.5	48,900	705	1278	571 704	-7. 8	51,200
543	803 1259	655 1143	-16.2 -8.0	45,900 25,200	626 627	1096 942	1089 548	-10.2 -13.3	27,200 53,000	706 707	1841 1018	1386	-0.7 -11.7	43,300 16,900
544 545	856	1526	-15.0	12,200	628	809	621	-16.0	48,000	709	1074	1145	-10.7	25,100
546	803	1071	-16.2	27,800	629	899	979	-14.1	31,300	710	293	889	<-35.0	34,800
547	1162	274	-9.3	98,400	630 631	1135 979	1321 615	-9.6	19,100 48,300	712 713	720 1386	412 841	-18.5 -6.4	66,600 36,800
548 549	128 1355	1321 1122	<-35.0 -6.8	19,000 25,900	632	1542	1076	-12.5 -4.1	27,600	714	1328	263	-7.1	103,100
550	595	866	-23.0	35,800	633	1345	814	-6.9	38,000	715	698	433	-19.1	63,900
552	1369	494	-6.6	57,500	634	409	950	-32.2	32,400	716	701	481	-19.0	58,700
553	992	405	-12.2 -9.8	67,600 66,900	635 636	1165 774	704 604	-9.2 -17.0	43,300 49,000	717 718	1875 575	699 702	-0.5 -23.9	43,600 43,400
555 5 5 6	1125 705	410 975	-18.9	31,400	637	1263	524	-8.0	54,800	719	1216	204	-8.6	140,400
557	1477	1030	-4.9	29,300	638	952	411	-13.1	66,700	721	1069	464	-10.8	60,400
558	980	583	-12.5	50,400	639	1717	575	-2.1	51,000	722 723	1272 958	506 822	-7.9 -13.0	56,400
559	700	1109 621	-19.1 -11.5	26,400 48,000	640 641	994 165	292 1224	-12.1 <-35.0	92,000 22,400	723	763	395	-17.3	37,700 69,100
560 562	1028 898	794	-14.1	38,900	642	803	251	-16.2	108,900	725	720	916	-18.5	33,700
564	.789	1446	-16.6	14,900	643	719	296	-18.5	90,700	726	1476	415	4.9	66,200
565	777	766	-16.9	40,200	644	1100	294	-10.2	91,400 21,000	727 728	1846 510	473 783	-0.7 -27.3	59,400 39,400
566	980	328 611	-12.5 -4.4	81,900 48,600	645 646	534 1153	1263 1038	-26.1 -9.4	29,000	729	1217	1126	-8.6	25,800
567 569	1519 1212	661	-8.6	45,600	648	1246	204	-8.2	140,000	730	1858	724	-0.6	42,300
570	760	594	-17.4	49,700	649	14	1406	<-35.0	16,200	731	665	765	-20.2	40,300
571	618	956	-21.9	32,100	650 651	1713 1986	1049 1183	-2.1 >0.0	28,600 23,800	733 734	1321 719	312 427	-7.2 -18.5	85,900 64,600
573	1142 532	771 787	-9.6 -26.2	40, 000 39,300	652	1378	816	-6.5	38,000	735	1101	473	-10.2	59,500
574 575	771	250	-17.1	109,200	653	1442	1165	-5.5	24,400	736	1359	569	-6.7	51,400
576	1068	534	-10.8	54,100	654	650	806	-20.8	38,400	738	696	220	-19.2	127,600
577	822	734	-15.7	41,800	655 656	1111 1095	551 861	-10.0 -10.3	52,700 36,000	739 740	687 1205	409 256	-19.5 -8.7	67,000 106,200
578	914 1064	754 794	-13.8 -10.8	40,800 38,900	657	1524	540	-10.3	53,600	741	995	563	-12.1	51,900
579 580	1524	714	-4.4	42,800	658	1777	860	-1.4	36,000	742	898	596		49,500
581	1392	783	-6.3	39,400	659	391	584	-33.4	50,400	743	881	181	-14.5	165,900
582	982	686	-12.4	44,200	660	977	565	-12.5	51,700	744 745	1951 726	686 168		44,200 183,600
584 585	1487 758	672 731	-4.8 -17.4	45,000 41,900	661 662	658 732	166 312	-20.5 -18.1	187,500 . 86,100	745	999	643		46,600
585 586	687	1152	-19.5	24,900	663	1787	567	-1.2	51,500	748	182	1503	<-35.0	13,000
587	930	523	-13.5	55,000	664	888	268	-14.4	100,900	749	2005	649		46,300 51,000
588	1888	774	-0.4	39,900 58 300	665 666	889 715	775 221	-14.3 -18.6	39,800 126,300	. 750 751	1448 792	575 266		51,000 101,900
589 590	642 1317	485 519	-21.1 -7.3	58,300 55,300	666 667	781	227	-16.8	122,400	752		296		90,600
590 591	65	1548	<-35.0	11,500	668	646	165	-21.0	189,100	754	664	254	-20.3	107,000
592	1014	614	-11.7	48,400	669	1116	353	-9.9	76,300	755	1195	184		161,000
593	732	176	-18.1	172,300	670 671	1382	643 780	-6.4 -25.3	46,600 39,200	756 757	1821 909	1113 246		26,300 111,000
594 506	1627 1009	478 1426	-3.0 -11.8	59,000 15.500	671 673	547 984	789 746	-25.3 -12.4	41.200	760		133		264.900
595	1009	1420	-11.0	. 3.550	0.0		,0							

MSN	X	Y	CPKol	SDSMW	MSN	х	Y	СРКоІ	SDSMW	MSN	X	Y	CPKol	SDSMW
			-6.2	41,800	648	1863	271	-0.6	99,500	939	1197	827	-8.8	37,500
761	1399	733 1085	-6.2 -5.9	27,300	849	1166	523	-9.2	54,900	941	1765	885	-1.5 ~~ 7	35,000
763 764	1416 2020	569	>0.0	51,400	850	1535	1024	-4.2	29, 600 37, 500	942 943	602 312	472 498	-22.7 <-35.0	59,600 57,100
765	651	475	-20.8	59,300	851	1035	826 542	-11.4 -15.5	53,400	944	993	491	-12.1	57,700
766	1052	1149	-11.1	25,000 59,900	852 855	634 499	220	-13.3 -27.8	127,100	945	1300	269	-7.5	100,300
767	1968	468	>0.0 -7.1	59,900 44,300	856	1063	194	-10.9	150,500	946	630	423	-21.6	65,100
768	1330 1970	685 613	>0.0	48,500	857	887	890	-14.4	34,800	947	187	736 344	<-35.0 -6.5	41,600 78,200
769 770	857	617	-15.0	48,200	858	1448	639	-5.4	46,900 86,200	948 949	1380 1766	665	-1.5	45,400
771	1337	974	-7.0	31,500	859	706 1070	311 1066	-18.9 -10.7	28,000	950	1038	193	-11.3	151,000
773	1576	502	-3.7 -12.8	56,700 37,600	860 861	472	347	-28.8	77,600	951	860	152	-14.9	213,000
775	969	824 708	-5.5	43,100	862	674	480	-19.9	58,800	952	957	701	-13.0	43,400
776 777	1438 1539	458	4.2	61,000	864	1307	499	-7.4	57,000	954 955	503 1938	547 712	•27.6 >0.0	53,000 42,900
778	850	434	-15.1	63,800	865	645	887	-21.0 -15.6	34, 900 30, 300	955 957	1010	816	-11.8	37,900
779	700	411	-19.1	66,800 25,500	866 868	827 685	1004 494	-19.5	57,400	959	768	174	-17.2	174,900
780	1052	1136	-11.1 -6.0	25,500 54,400	869	1807	402	-1.0	68,000	960	596	419	-23.0	65,700
784	1413 1364	529 885	-6.7	35,000	670	1323	783	-7.2	39,400	961	557	409	-24.8	67,100 83,900
785 786	1822	835	-0.9	37,100	871	1228	1031	-8.4	29,300	962 963	887 564	320 334	-14.4 -24.5	80,500
787	893	-392	-14.3	69,500	872	1904	346	-0.3	77,700 46,400	964	969	1155	-12.8	24,800
790	616	882	-22.0	35,100	873 874	556 1540	647 756	-24.8 -4.2	40,700	965	671	255	-20.0	106,600
791	451	1429	-29.8 -16.9	15,400 72,000	875	1566	777	-3.8	39,700	966	1204	798	-8.7	38,700
792	777 1536	377 1543	-4.2	11,700	876	1198	351	-8.8	76,800	967	910	154	-13.9	210,300
793 794	1461	807	-5.1	38,300	877	1076	720	-10.6	42,500	968 969	609 1285	1048 206	-22.3 -7.7	28,700 138,900
796	388	546	-33.6	53,100	878	1161	1111	-9.3 20.0	26,400 40,700	970	822	232	-15.8	119,300
797	1126	212	-9.8	133,700	879 880	647 1756	757 594	-20.9 -1.6	49,700	971	976	437	-12.6	63,400
798	933	437 593	-13.5 -5.9	63,400 49,800	881	1543	278	-4.1	97,100	972	403	567	-32.6	51,600
799 800	1420 1759	279	-1.6	96,500	883	1432	890	-5.7	34,800	974	279	495	<-35.0 -15.3	57,400 31,200
801	624	865	-21.7	35,800	884	922	689	-13.7	44,100 66,400	975 976	844 1124	981 295	-15.3 -9.8	91,100
802	898	547		53,000	885 686	1103 1501	414 607	-10.1 -4.6	48,900	977	994	664	-12.1	45,400
803	1775	1468 196	-1.4 -24.0	14,200 148,400	887	798	1103	-16.3	26,600	978	1612	642	-3.2	45,700
804 805	573 203	494	<-35.0	57,400	888	636	634	-21.3	47,200	979	749	1141	-17.7	25,300 46,700
806	980	1039		29,000	889	951	759	-13.1	40,600	980 981	1064 1197	642 911	-10.8 -8.8	33,900
807	902	308		87,200	890		548 229	-18.6 -9.8	52,900 121,200	983	1762	1508		12,800
808	625	827		37,500 29,900	891 892	1123 891	413	-14.3	66,400	984	1344	317		84,700
809	1851 440	1015 573		51,100	894	1245	234	-8.2	117,800	985		1105		26,600
810 811	1358	249		109,700	895		346	>0.0	77,700	987		1159 555		24,600 52,400
812		393	-15.1	69,400	896		626	-7.2	47,700 51,300	988 990		361		74,900
813	745	1246		21,600	897 898		570 428	-31.4 -20.3	64,500	991		317		84,500
814		810		38,200 46,500	899		243		113,000	992		928		33,300
815 816		645 313		85,700	900		703	-21.7	43,400	993		701		
817				24,000	901		1094		27,000	994 995		811 461		60,700
818	1771	790		39,100	903 904		229 520		121,000 55,200	996		847		36,600
819		263		103,100 74,600	905		889		34,800	997	7 1815	579		
820 821		362 279			907		824	-14.4	37,600	998				
822				139,200	908		1303		19,700	999 1000				
823	1517	654			910		1544 301		11,700 89,1 0 0	100				
824					91° 91°					100			B -1.9	58,900
825					91	_		_	44,100	100				
826 827					910	1442				100				
828			5 -13.4		91					100 100				
830					91: 92:			_		101				
831					92					101		54		
832			.		92				113,200	101				
833 834				50,500	92	4 1131	318			101				
837		_	B -17.6	41,100	92					101 101				
838	635	83			92					101			_	
839					92 92					101	7 172	2 42	4 -2.0	0 65,000
840					92				38,000	101				
841 842	-	-			93	1 1609	67			102				
843			9 -7.2	46,300	93					102 102			34 -16. 33 -9.	
84	4 1727	7 30			93					102				9 84,600
84					93 93					102	24 78	5 44	46 -16 .	7 62,400
840			-		93	-					25 129	0 7:	39 -7.	7 41,500
841	7 673	, 120	J 10.0			_								

ASN	×	Y	CPKol	SDSMW	MSN	x	Y	CPKpl	SDSMW	MSN	×	Y	CPKol	SDSMW
1026	405	552	-32.5	52,600	1153	921	1158	-13.7	24,700	1246	547	577	-25.3	50,800
1027	1298	848	-7.5	36,500	1154	1594	864	-3.5	35,900	1247 1249	530 516	576 572	-26.3 -27.0	50,900 51,200
1028	856	547	-15.0	53, 000 123, 200	1161 1162	637 623	400 397	-21.3 -21.8	68,400 68,800	1250	973	536	-27.0 -12.7	53,900
1030	1284	226 822	-7.7 -12.3	37,700	1163	665	397	-20.2	68,700	1251	607	532	-22.4	54,200
1031	986 1547	403	-4.1	67,900	1168	564	528	-24.4	54,500	1252	665	529	-20.2	54,400
1032 1033	1381	551	-6.4	52,700	1170	552	529	-25.0	54,500	1253	899	766	-14.1	40,200
1034	1525	496	-4.3	57,200	1171	538	524	-25.9	54,800	1254	1311	746	-7.4	41,200
1035	1128	645	-9.7	46,500	1172	545	514	-25.5	55,700	1255	1300	761 712	-7.5 0.0	40,400 42,900
1036	1226	274	-8.5	98,300	1174 1176	1099 1304	522 586	-10.2 -7.5	55,000 50,200	1257 1258	1938 1806	718	-1.0	42,600
1039	1761	262 839	-1.6 -25.7	103,600 36,900	1177	1366	539	-6.6	53,700	1259	1727	715	-2.0	42,700
1040	541 818	910	-15.8	34,000	1178	1608	702	-3.3	43,400	1260	1629	713	-3.0	42,800
1041 1044	1036	485	-11.3	58,300	1179	1485	224	-4.8	124,900	1261	1555	717	-4.0	42,600
1045	1439	407	-5.5	67,300	1180	1459	224	-5.2	124,900	1262	1468	717	-5.0	42,600
1047	1540	250	-4.2	109,200	1181	1431	223	-5.7	125,100	1263	1413	722	-6.0 7.0	42,400 42,600
1048	1576	635	-3.7	47,100 66,700	1182	1407	223 224	-6.1	125,200 124,700	1264 1265	1340 1263	717 717	-7.0 -8.0	42,600
1049	1089	411	-10.4 -13.2	66,700 28,900	1183 1184	1383 1454	182	-6.4 -5.3	164,400	1265	1182	720	-9.0	42,500
1050	949	1040 818	-13.2	37,800	1185	1422	183	-5.8	162,600	1267	1110	717	-10.0	42,600
1051 1052	426 1583	1385	-3.6	16,900	1186	1394	182	-6.3	164,300	1268	1055	717	-11.0	42,600
1053	779	1092	-16.8	27,000	1189	1171	214	-9.2	131,800	1269	999	717	-12.0	42,600
1054	1613	620	-3.2	48,000	1190	1457	286	-5.2	94,200	1270	959	715	-13.0	42,700
1055	1380	377	-6.5	72,000	1191	686	1114	-19.5	26,200	1271	905	712	-14.0	42,900
1056	284	663	<-35.0	45,500	1192	265 403	893 1292	<-35.0	34,700 20,000	1272 1273	857 810	714 705	-15.0 -16.0	42,800 43,300
1058	1261	746	-8.0 -33.3	41,200 49,000	1193 1194	344	1275	-32.6 <-35.0	20,600	1274	774	711	•17.0	42,900
1060	393 1817	605 645	-33.3 -0.9	46,600	1195	505	1311	·27.6	19,400	1277	737	708	-18.0	43,100
1061 1062	1245	746	-8.2	41,200	1196	572	1293	-24.1	20,000	1278	702	711	-19.0	42,900
1064	1258	792	-8.1	39,000	1197	639	1502	-21.2	13,000	1279	671	710	-20.0	43,000
1065	705	934	-18.9	33,000	1198	637	1402	-21.3	16,300	1280	645	710	-21.0	43,000
1066	1181	734	-9.0	41,800	1199	614	1407	-22.1	16,200	1281	617	707	-22.0	43,100
1067	529	658	-26.3	45,800	1200	637	1431	-21.3	15,400 16,600	1282 1283	595 573	704 700	-23.0 -24.0	43,300 43,500
1068	508	696 604	-27.4 -0.3	43,700 49,100	1201 1202	1095 1719	1394 1545	-10.3 -2.1	11,600	1284	552	695	-25.0	43,700
1069 1071	1898 873	609	-14.7	48,700	1203	791	668	-16.5	45,200	1285	536	694	-26.0	43,800
1073	1768	1128	-1.5	25,800	1204	964	1021	-12.9	29,700	1286	515	687	-27.0	44,200
1075	836	773	-15.4	39,900	1205	313	195	<-35.0	148,700	1287	496	683	-28.0	44,400
1076	1863	861	-0.6	36,000	1208	306	194	<-35.0	149,800	1288 1289	467 447	669 667	-29.0 -30.9	45,200 45,300
1078	826	566	-15.7 -12.7	51,600 58,500	1209 1210	320 326	197 197	<-35.0 <-35.0	147,400 146,600	1290	427	655		45,900
1081	971	483 202	-12.7	142,300	1211	394	294	-33.2	91,400	1291	412	655		45,900
1083 1085	1697 1157	794	-9.4	38,900	1212	402	294	-32.7	91,200	1292	397	652		46,100
1090	620	910	-21.9	34,000	1214	386	294	-33.7	91,400	1293	381	654		46,000
1092	1867	597	-0.5	49,500	1215	641	329	-21.2	81,600	1294	365	653		46,100
1093	2019	894	>0.0	34,600	1216	660	329	-20.4	81,600	1295	348	653	<-35.0	46,100
1094	1546	538	-4.1	53,700	1217	914 873	266 245	-13.8 -14.7	101,800 112,000					
1095	1545	477	-4.1 <-35.0	59,100 33,000	1218 1219	970	372	-12.7	72,900					
1098 1099	61 1954	935 237	>0.0	116,000	1220	1021	298	-11.6	90,100					
1101	588	1048	-23.3	28,600	1221	1392	205	-6.3	139,500					
1102	1050	667	-11.1	45,200	1222	1354	203	-6.8	141,800					
1103	457	797	-29.5	38,800	1223	1362	205	-6.7	139,500					
1105	1884	532	-0.4	54,200	1224	673	540	-19.9	53,600 53,400					
1106	1714	649	-2.1	46,300 53,100	1225 1226	614 603	542 539	-22.1 -22.6	53,600					
1107	1717	546 722	-2.1 >0.0	42,400	1227	696	623	-19.2	47,800					
1108 1111	1976 547	1066	-25.3	28,000	1228	. 707	628		47,500					
1112	1348	621	-6.9	48,000	1229	475	447		62,300					
1115	1385	762	-6.4	40,400	1230	466	1282		20,400					
1116	1078	816	-10.6	38,000	1231	759	1461	-17.4	14,400					
1117	975	787	-12.6	39,300	1232	1324	1170		24,200					
1118	1202	933	-8.7	33,100	1233 1234	1583 1865	1005 809		30,300 38,200					
1119	1022	1076	-11.6 -0.3	27,600 48,300	1234	1812	817		37,900					
1120	1905 1512	616 1301	-4.5	19,700	1236	1411	703		43,400					
1121 1122	1114	677	-9.9	44,700	1237	1392	682		44,500					
1123	1464	452	-5.1	61,700	1238	794	410		66,900					
1125	1048	857	-11.1	36,200	1239	769	407		67,300					
1126	1122	802	-9.8	38,600	1240	740	406		67,500					
1128	1722	892	-2.1	34,700	1241	743	511		55,900 56,000					
1133	1098	825	-10.2	37,500 51,400	1242	713 682	510 509		56,000 56,100					
1139	1830	569 1182	-0.8 -17.3	51,400 23,800	1243 1244	663	509		56,500					
1147	764													

Table 2. Table of some identified proteins	proteins	MCNI'S	Basis for identification
POP name	Protein name		
HOUH WHO IS C. SOI	-loldondin-biorotavanbad-s-6	137, 159 F	Pure protein and antibody provided by Dr. T.M.
	dehydrogenase, an enzyme of		Penning, Department of Pharmacology, School
	steroid metabolism		of Madicine, University of Pennsylvaria.
IDS:ACTIN_BETA	β cellular actin, a cytoskeletai protein	a.	
IDS:ACTIN_GAMMA	y cellular actin, a cytoskeletal protein	1 68	Homologous position with respect to other mammallan
		•	systems Orodominanto in rational
IDS:AFBOWING	Apo A-I plasma lipoprotein, mature form	236, 463	Presence in rat plasma, regulation by some lipid-
	(tentative).		lowering drugs
IDS:CALMODULIN	Calmodulin, an acidic cytosolic calcium-	123, 649	Homologous position with respect to other mammatian
IDS:CATALASE	Catalase (peroxisomal)	54, 61, 106	Presence in purified peroxisomes, similarity in position
	Second MOO and the first second second	1267 1206	to mouse catalase
IDS.CFASFOIS	opois controlled by the OTA charge standards (not rat liver proteins)		
IDS:CPS	Carbamoyl phosphate synthase	114, 157, 167, 174, 1184, 1185, 1186, 1222	Pure protein provided by Dr. Margaret Marshall, Department of Pharmacology, Medical School.
IDS:CYTOCHROME_B5	Cytochrome b5	87, 477	Pure protein provided by Ur. Andrew Parkinson, Department of Phermacology, Toxicology and Therapeutics, University of Kansas Medical
	ciatora sciboid bine standardi	756	Center Pure protein provided by Dr. Nathan Bass, Department
IDS:rABP·L	Liver fairy-acto binoing protein		of Medicine, University of California School of Medicine. San Francisco
IDS:HMG-COA_SYNTHASE	Cytosolic HMG-CoA Synthase	133, 144, 235, 413	Antibody provided by Dr. Michael Greenspan, Merck Sharp & Dohme Research Laboratories,
		415 734	Rahway, NJ Homologous position with respect to other mammallan
IDS:LAMIN_B	Lamin D, a mucieal profess		systems
IDS:MITCON:1	Mitcon:1 (F1 ATPase B subunit), a	17, 49, 71, 340, 1245, 1246, 1247, 1249	Homologous position with respect to other mammalian systems, presence in mitochondria
S-MUTCON-S	Mitcon: 2. a mitochondrial matrix stress	15, 25, 110, 1241, 1242, 1243, 1244	Homologous position with respect to other mammallan
#:NOO 18:50	protein equivalent to E.	18 35 326 600 1238 1239 1240	systems, presence in miloculouse Homologous position with respect to other mammalian
IDS:MITCON:3	Mitcon:3, a mitochondrial matrix stress	16, 33, 226, 600, 1236, 1235, 1245	systems, presence in mitochondria
IDS:NADPH_P450_RED	NADPH cytochrome P-450 reductase, frequently co-induced with P-450's	175, 251, 812	Pure protein provided by Ur. Andrew Parkinson, Department of Pharmacology, Toxicology and Therapeutics, University of Kansas Medical
		221 121 121	Center Sequence Information obtained by R.M. Van Frank,
IDS:PDI	Protein disulphide (somerase 1	108, 1170, 1171, 1172	Lilly Research Laboratories, Indianapolis
IDS:PLASMA_PROTEINS	Rat plasma proteins observed in liver	21, 28, 33, 44, 72, 102, 115, 197, 236, 246, 246, 257, 293, 332, 347, 364, 369, 419, 432, 463, 468, 518, 562, 605, 623, 666, 667, 725,	Plasma coelectrophorests studies
IDS-PRO. AT BLIMIN	Serum albumin precursor	738, 790, 865, 903, 928 47, 93	Relative position to mature albumin, presence in micro-
		470 4400 4481 4482 4183	Paviica, R.J., et al., BBA (1990) <i>1022</i> 115-125.
IDS:PYRCARBOX IDS:SOD	Pyruvate carboxylase Superoxide dismutase	135	Sequence information obtained by R.M. Van Frank, Lilly Research Laboratories, Indianapolis
IDS:TUBULIN ALPHA	a tubulin, a cytoskeletal protein	56, 132, 1224, 1252	Homologous position with respect to other mammalian systems
ATAN IN BETA	8 tubulin, a cytoskeletał protein	50, 1225, 1226, 1251	Homologous position with respect to other mammallan
			SAIGHE

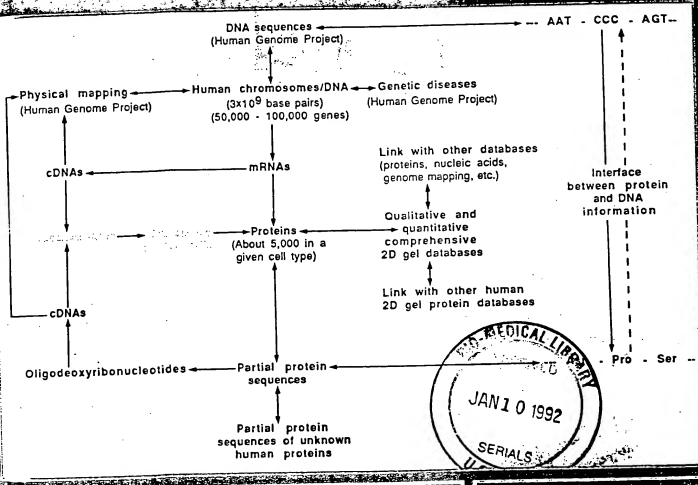
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Table 3. Computed p/s of two sets of carbamylated protein standards: Rabbit muscle CPK and human hemoglobin (Hb)

	emoglobin (nb)	PIR	#ASP	#GLU	#HIS	#LYS	#ARG	NH2-	Calc	Real
	Protein Name	Name	3.9	4.1	6.0	10.8	12.5	7.0	pl	СРК
0	Rabbit muscle CPK	KIRBCM	28	27	17	34	18	1	6.84	0.0
-1	•		28	27	17	33	18	1	6.67	-1
-2			28	27	17	32	18	. 1	6.54	-2
-3			28	27	17	31	18	1	6.42	-3
-4			28	27	17	30	18	1	6.31	4
-5			28	27	17	29	18	1	6.21	-5 -6
-6			28 28	27	17	28	18	1	6.12 6.03	-0 -7
-7			28	27 27	17 17	27 26	18 18	1 1	5.94	-7 -8
-8			28	27	17	25	18	1	5.85	-9
-9 -10			28	27	17	24	18	i	5.76	-10
-10			28	27	17	23	18	i	5.67	-11
-12			28	27	17	22	18	1	5.58	-12
-13			28	27	17	21	18	1	5.48	-13
-14			28	27	17	20	18	1	5.39	-14
-15			28	27	17	19	18	1	5.29	-15
-16			28	27	17	18	18	1	5.20	-16
-17			28	27	17	17	18	1	5.12	-17
-18			28	27	17	16	18	1	5.04	-18
-19			28	27	17	15		1	4.96	-19
-20			28	27	17	14	18	1	4.89	-20
-21		•	28	27	- 17	13	18	1	4.83	-21
-22			28	27	17	12	18	1	4.77	-22
-23			28 28	27 27	17 17	11 10	18 18	1	4.71 4.66	-23 -24
-24			28	27	17	9			4.61	-25
-25 -26			28	27	17	8		i	4.56	-26
-20 -27			28	27	17	7		i	4.52	-27
-28			28	27	17	6		1	4.48	-28
-29			28	27	17	5		1	4.44	-29
-30			28	27	17	4	18	1	4.40	-30
-31			28	27	17	3	18	1	4.36	-31
-32			28	27	17	2	18	1	4.32	-32
-33			28	27	17	1		1	4.29	-33
-34			28	27	17	0		1	4.25	-34
-35			28	27	17	0	18	0	4.22	-35
0	Hb-beta, human	HBHU	7	8	9	. 11	3	1	7.18	
-1			7	8	9	. 10		1	6.79	
-2			.7	8	9	9		1	6.53	-1.8
-3			7	8	9	8		1	6.32	-3.2
-4			7	8	9	7		1	6.13	-5.3
-5	•		7	8	9	6	3	1	5.96	-7.2 10.0
-6	÷		7 7	8	9	5 4		1	5.78 5.59	-10.0 -12.3
-7 .			7	8 8	9	3	3	1	5.39 5.37	-12.3 -15.5
-8 -			7	8	9	2	3	1	5.14	-13.3
-9			7	8	9	1		1	4.91	-21.0
-10			7	8	9	Ö		1	4.71	-25.5
-11 -12			7	8	9	Ö		Ö	4.54	-27.2
-12										

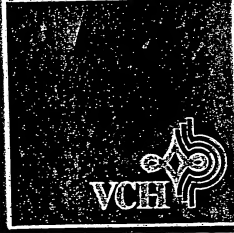
Table 4. Computed p/s of some known proteins related to measured CPK p/s

-	Protein Name	PIR Name	#ASP 3.9	#GLU 4.1	#HIS 6.0	#LYS 10.8	#ARG 12.5	Calc	Real CPK
		KIRBCM	28	27	17	34	18	6.84	0.0
0	Creatine phospho kinase (CPK), rabbit muscle	FZRTL	5	13	2	16	2	7.83	-3.0
1	Fatty acid-binding protein, rat hepatic	MGHUB2	7	8	4	8	5	6.09	-5.0
2	b2-microglobulin, human	SYRTCA	72	96	28	95	56	5.97	-5.5
3	Carbamoyl-phosphate synthase, rat	ABRTS	32		15	53	27	5.98	-6.2
4	Proalbumin (serum albumin precursor), rat	ABRTS	32		15		24	5.71	-9.0
5	Serum albumin, rat	A26810	8		10	9	4	5.91	-9.2
6	Superoxid dismutase (Cu-Zn, SOD), rat	A28807	34		9	49	21	5.92	-9.2
7	Phospholipase C, phophoinositide-specific (?), rat	ABHUS	36		16	60	24	5.70	-11.9
8	Albumin, human	A24700	18				12	5.32	-13.7
9	Apo A-1 lipoprotein, rat	LPHUA1	16				17	5.35	-14.3
10	proApo A-I lipoprotein, human	RDRTO4				_	36	5.07	-15.6
11	NADPH cytochrome P-450 reductase, rat	VAHU	18					5.04	-16.9
12	Retinol binding protein, human	ATRTC	23					5.06	-17.2
13	Actin beta, rat	ATRTC	20					5.07	-16.8
14	Actin gamma, rat	LPHUA1	16					5.10	-17.5
15	Apo A-I lipoprotein, human	LPHUA4							-19.7
16	Apo A-IV lipoprotein, human	UBRTA	27					4.66	-19.8
17	Tubulin alpha, rat	PWBOB				22			
18	F1ATPase beta, bovine	UBPGB							
19	Tubulin beta, pig	ISRTSS							
20	Protein disulphide isomerase (PDI), rat hepatic	CBRT5	11			6 10			
21	Cytochrome b5, rat								
22	Apo C-II lipoprotein, human	LPHUC2		-		`		,	
	Amino acid pl assumed in calulation:		3.	9 4.	1 6.	0 10.8	8 12.5	5	



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